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## BLOCK-BASED BAYESIAN EPISTASIS ASSOCIATION MAPPING WITH APPLICATION TO WTCCC TYPE 1 DIABETES DATA<sup>1</sup>

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Interactions among multiple genes across the genome may contribute to the risks of many complex human diseases. Whole-genome single nucleotide polymorphisms (SNPs) data collected for many thousands of SNP markers from thousands of individuals under the case–control design promise to shed light on our understanding of such interactions. However, nearby SNPs are highly correlated due to linkage disequilibrium (LD) and the number of possible interactions is too large for exhaustive evaluation. We propose a novel Bayesian method for simultaneously partitioning SNPs into LD-blocks and selecting SNPs within blocks that are associated with the disease, either individually or interactively with other SNPs. When applied to homogeneous population data, the method gives posterior probabilities for LD-block boundaries, which not only result in accurate block partitions of SNPs, but also provide measures of partition uncertainty. When applied to case–control data for association mapping, the method implicitly filters out SNP associations created merely by LD with disease loci within the same blocks. Simulation study showed that this approach is more powerful in detecting multi-locus associations than other methods we tested, including one of ours. When applied to the WTCCC type 1 diabetes data, the method identified many previously known T1D associated genes, including *PTPN22*, *CTLA4*, *MHC*, and *IL2RA*. The method also revealed some interesting two-way associations that are undetected by single SNP methods. Most of the significant associations are located within the MHC region. Our analysis showed that the MHC SNPs form long-distance joint associations over several known recombination hotspots. By controlling the haplotypes of the MHC class II region, we identified additional associations in both MHC class I (*HLA-A*, *HLA-B*) and class III regions (*BAT1*). We also observed significant interactions between genes *PRSS16*, *ZNF184* in the extended MHC region and the MHC class II genes. The proposed method can be broadly applied to the classification problem with correlated discrete covariates.

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**1. Introduction.** A recent genome-wide association (GWA) study of 14,000 cases of seven human genetic diseases and 3,000 shared controls by the Wellcome Trust Case Control Consortium [WTCCC (2007)] represents a thorough validation of the GWA approach. By testing hundreds of thousands of single nucleotide polymorphisms (SNPs) in the human population (Affymetrix 500k SNP), the study has identified many SNPs associated with seven complex diseases [WTCCC (2007)]. Each SNP consists of two *alleles* taking values 0 or 1, and there are three possible combinations of the two alleles: (0, 0), (0, 1), (1, 1), disregarding the order. Each combination is called a *genotype*, representing wild homozygote, heterozygote, and mutant homozygote, respectively. In a case-control study, a SNP is said to be associated with the disease if the genotype (or allele) distribution at the SNP is different between cases and controls. In addition to testing individual SNPs, it has also been anticipated that epistatic interactions among SNPs, defined as multiple SNPs jointly associated with the disease, may be responsible for significantly elevating the risks of some human complex diseases. Due to computational and methodological limitations, however, efforts on detecting disease-related epistatic interactions among SNPs in the WTCCC study have been limited.

In the past few years, many approaches have been developed for case-control studies to detect epistasis associations. Most methods cannot be applied to GWA studies due to their computational limitations except for some recently developed methods, such as the stepwise logistic regression method [Marchini, Donnelly and Cardon (2005)] and the Bayesian epistasis association mapping (BEAM) algorithm [Zhang and Liu (2007)]. It has been demonstrated that BEAM is capable of detecting high-order interactions in GWA studies and is more powerful than other existing methods [Zhang and Liu (2007)]. A limitation of BEAM, however, is its model assumption that a Markov chain can capture the dependence structure of the SNPs in the data. It is well known that linkage disequilibrium (LD) between adjacent SNPs exhibits block-wise structure in the human genome [The International HapMap Consortium (2005), Reich et al. (2001)]. SNPs within blocks are highly correlated and the correlation is broken down by recombination events at block boundaries. A simple Markov model cannot capture this important block structure when analyzing dense SNPs.

Previous studies have shown that using *haplotypes*, defined as allele combinations over multiple nearby SNPs inherited from one of the parents, cannot only reduce the high computation cost in GWA studies, but also improve the detection power of association mapping [Kuno et al. (2004); Zöllner and Pritchard (2005); Johnson et al. (2001); Zhang et al. (2002a); de Bakker et al. (2005)]. In particular, Nielsen et al. (2004) showed that when moderate to high levels of LD exist, haplotype tests tend to be substantially more powerful. Kuno et al. (2004) demonstrated in a real disease study that single-SNP tests were not significant even at SNP loci close to the mutation site (APRT\*J), whereas the haplotype block data yielded sufficient statistical significance.

Similar to haplotypes, we define *diplotypes* as genotype combinations over multiple nearby SNPs. Diplotypes are directly observed in GWA studies, whereas haplotypes have to be inferred using computationally expensive algorithms. Throughout the paper, we discuss our method and results on diplotype associations with the disease, although the method is directly applicable to haplotype data. Our focus on diplotype association is mainly due to the computational concern, where inferring unobserved haplotypes will be extremely time consuming. It is also possible that testing diplotype associations could be more powerful than testing haplotype associations, depending on the underlying disease model. On the other hand, if haplotype associations are of the interests, users can first infer haplotypes using an available haplotype inference algorithm, and then input the inferred haplotypes into our method. The degrees of freedom of our model will automatically accommodate the different inputs.

In this paper we extend the BEAM model to address the block structures in the human genome. We refer to SNP block structures as LD-blocks. By partitioning SNPs into LD-blocks, a naïve extension of BEAM is to treat each LD-block as a genetic marker, with diplotypes in the block being treated as alleles. This approach, however, may not be optimal for association mapping. First, criteria utilized in existing block-partitioning algorithms do not directly aim at optimizing the power of association mapping. Second, many regions in the human genome demonstrate vague structural patterns, of which a measure of uncertainty in block structures should be provided. Simulation studies have shown that LD-block structures can be affected by marker density [Wang et al. (2002); Phillips et al. (2003); Wall and Pritchard (2003)], population structure [Wang et al. (2002); Stumpf and Goldstein (2003); Zhang et al. (2003); Anderson and Slatkin (2004)], and gene conversion [Przeworski and Wall (2001)].

We propose a Bayesian model to simultaneously infer LD-blocks and select SNPs within blocks for disease association mapping. The model partitions the genome into discrete blocks, within which the diversity of diplotypes is limited. Block structures are iteratively updated such that disease associations are detected and summarized from a variety of likely block partitions. This approach takes into account the uncertainty in block structures and optimizes the detection power by searching for the best partitions around disease-associated SNPs. Our method detects combinations of SNPs within and between blocks for marginal and epistatic associations to the disease status. Using LD-blocks, our method also automatically filters out artificial associations created merely by LD with nearby authentically associated SNPs. We show that the new method, BEAM2, is more powerful than the original BEAM (renamed BEAM1 henceforth).

By applying BEAM2 to the type 1 diabetes (T1D) data from WTCCC (2007), we obtained all the previously identified single SNP associations in the WTCCC T1D data. We further observed some interesting two-way joint associations not detectable by single-SNP methods. The strongest T1D associations occur in the

well-known Major Histocompatibility Complex (MHC) region, in which we observed long-distance joint association patterns over several millions of base pairs (Mb). Since this pattern may be partially caused by the extended LD from the MHC class II region, we further controlled the structure of MHC class II using a logistic regression model, and tested additional effects of SNPs over the extended MHC region as well as the MHC class I and class III regions. We observed strong associations in the MHC class I and class III regions, and found significant interaction associations between genes *PRSS16*, *ZNF184* in the extended MHC region and the MHC class II genes.

The article is organized as follows. In Section 2 we first introduce a LD-block model for the genotype distribution among multiple SNPs. We model the genotype distribution within a block of SNPs by multinomial distributions and assume that the joint distribution of all SNPs is the product of the distributions of individual blocks (i.e., assuming block independence). In Section 3 we extend the LD-block model to incorporate disease associated SNPs and epistasis and describe Monte Carlo Markov chain (MCMC) algorithms to make inference from the joint model for both LD-blocks and disease associations. In Section 4 we briefly review our previously developed Bayes factor-based test statistic, which is used to further evaluate the statistical significance of candidate epistasis detected by BEAM2, and discuss extensions of BEAM and BEAM2 for general classification problems. In Section 5 we demonstrate the superior performance of BEAM2 by simulation studies and real data applications. In Section 6 we report our results from applying BEAM2 to the T2D data from WTCCC (2007). We conclude the article with a short discussion in Section 7. More implementation details can be found in Supplemental Material [Zhang, Zhang and Liu (2011)].

**2. A Bayesian model for LD-block inference.** The data of interest consist of genotypes at a total of  $L$  SNP markers (or  $L$  covariates, each taking on 3 possible values) observed in  $N_d$  case and  $N_u$  control individuals. Let  $D = (D_1, \dots, D_L)$  denote a  $N_d \times L$  matrix of case genotypes, and  $U = (U_1, \dots, U_L)$  denote a  $N_u \times L$  matrix of control genotypes, where  $D_i$  and  $U_i$  denote vectors of genotypes observed at SNP  $i$  across individuals.

**2.1. Bayesian LD-block model.** Here we introduce a Bayesian LD-block partition model without considering disease association. Hence, we treat cases ( $D$ ) and controls ( $U$ ) as coming from the same population and use the combined data ( $D, U$ ) to describe the model. A *diplotype* for an LD-block of SNPs is defined as a particular genotype combination of all the SNPs in that block. We seek to partition the  $L$  markers into  $B$  consecutive blocks, so that the number of observed diplotypes within each block is small (strong correlation), and correlation between SNPs in different blocks is weak. Block partitions can be quite ambiguous in many genomic regions and can vary across samples in details. Diplotype block structures

obtained from available software are often based on ad hoc criteria that neither result in a proper uncertainty measure nor optimize the association mapping power.

The block variable  $B$  in our model consists of  $L$  binary indicators corresponding to the  $L$  SNPs in the data. An indicator is equal to 1 if the corresponding SNP is the start position of a block, and 0 otherwise. As a result,  $B$  uniquely defines a partition of SNPs into consecutive blocks. For a diplotype block  $[a, b]$  consisting of SNPs  $a, \dots, b-1$ , we let  $(n_h, m_h)$  denote the counts of a particular *diplotype*  $h$  in cases and controls, respectively. There are  $3^{b-a}$  possible diplotypes in the block. We assume that the diplotype of each individual follows independently from a multinomial distribution with frequency parameters  $\{p_h\}$ , and  $\{p_h\}$  follows a Dirichlet prior distribution,  $\text{Dir}(\{\alpha_h\})$ , where  $\{\alpha_h\}$  denotes a hyper-parameter (i.e., pseudo-counts). More precisely, letting  $\{n_h + m_h\}$  denote the combined counts of diplotype  $h$  observed in cases and controls, we have

$$\Pr(\{n_h + m_h\}|\{p_h\}) = \prod_{h=1}^{3^{b-a}} p_h^{n_h+m_h} \quad \text{and} \quad \Pr(\{p_h\}) = \frac{\Gamma(\alpha_\bullet)}{\prod_{h=1}^{3^{b-a}} \Gamma(\alpha_h)} \prod_{h=1}^{3^{b-a}} p_h^{\alpha_h-1},$$

where the subscript ' $\bullet$ ' denotes the sum of values over all subscripts. We can integrate out  $\{p_h\}$  by

$$\begin{aligned} & \int \Pr(\{n_h + m_h\}|\{p_h\}) \Pr(\{p_h\}) d(\{p_h\}) \\ &= \frac{\Gamma(\alpha_\bullet)}{\prod_{h=1}^{d-1} \Gamma(\alpha_h)} \int \prod_{h=1}^{3^{b-a}} p_h^{n_h+m_h+\alpha_h-1} d(\{p_h\}). \end{aligned}$$

Noting that the integrand on the right-hand side is proportional to the density function of  $\text{Dir}(n_h + m_h + \alpha_h)$  up to its normalizing constant, we obtain the marginal probability of the combined data of block  $[a, b]$  as

$$\begin{aligned} (1) \quad & P(D_{[a,b]}, U_{[a,b]}|[a, b] \text{ forms a block}) \\ &= \left( \prod_{h=1}^{3^{b-a}} \frac{\Gamma(n_h + m_h + \alpha_h)}{\Gamma(\alpha_h)} \right) \frac{\Gamma(\alpha_\bullet)}{\Gamma(n_\bullet + m_\bullet + \alpha_\bullet)}. \end{aligned}$$

We set  $\alpha_h = \rho/3^{b-a}$  for diplotype  $h$ , and by default, we let  $\rho = 1.5$ . Based on the likelihood equivalence principle [Heckerman, Geiger and Chickering (1995)],  $\{\alpha_h\}$  is chosen to be inversely proportional to the total number of possible diplotypes in a block, and, hence, the sum  $\alpha_\bullet$  remains a constant over different block sizes. Note that formula (1) can also be used to model case data ( $D$ ) or control data ( $U$ ) only, which is simply done by letting  $m_h = 0$ , or  $n_h = 0$ , respectively, for all  $h$ . Further assuming independence between blocks (which is not entirely true, but serves as a good approximation), the probability function  $P(D, U|B)$  of genotypes of all blocks can be expressed as the product of individual block probabilities defined in formula (1).

**2.2. Bayesian inference of SNP association based on blocks.** A saturated test of disease association for a diplotype block of  $Mz$  consecutive SNPs involves  $3^M - 1$  free parameters. When  $M$  is moderately large ( $M > 3$ ), the power of the test becomes exceedingly low. We propose a Bayesian model of disease associations using only a subset of markers in the block. Here, we only discuss joint associations for SNPs within a diplotype block, and we will address epistatic interactions in the next section. It therefore suffices to describe our model for only one block.

Let  $\{M\}$  denote the set of  $M$  SNPs in the block. We assume that only a subset  $\{x\}$  of SNPs of size  $x$  (often is 0 or 1) are truly associated with the disease, and the SNPs in  $\{M\} \setminus \{x\}$  are not associated with the disease given  $\{x\}$ . Thus, the diplotypes of SNPs in  $\{x\}$  are distributed differently and hence are modeled by two different distributions for cases and controls, respectively. Conditional on the diplotypes of SNPs in  $\{x\}$ , the diplotypes of SNPs in  $\{M\} \setminus \{x\}$ , however, follow a common distribution between cases and controls. The joint probability of the block data can therefore be expressed as

$$(2) \quad P(D_{\{M\}}, U_{\{M\}}) = P(D_{\{x\}})P(U_{\{x\}})P(D_{\{M\} \setminus \{x\}}, U_{\{M\} \setminus \{x\}} | D_{\{x\}}, U_{\{x\}}).$$

Here,  $P(D_{\{x\}})$  and  $P(U_{\{x\}})$  are modeled by two independent multinomial-Dirichlet distributions specified in formula (1), treating all SNPs in  $\{x\}$  jointly as a block. To model  $P(D_{\{M\} \setminus \{x\}}, U_{\{M\} \setminus \{x\}} | D_{\{x\}}, U_{\{x\}})$ , we combine the diplotypes of SNPs in  $\{M\} \setminus \{x\}$  in cases and controls together. These diplotypes are not directly associated with the disease given  $\{x\}$ , and thus have the same conditional distributions between cases and controls. Conditional on each possible diplotype  $h$  of SNPs in  $\{x\}$ , we model the conditional diplotype distribution of SNPs in  $\{M\} \setminus \{x\}$  again by a multinomial-Dirichlet distribution. It is straightforward to derive the expression

$$P(D_{\{M\} \setminus \{x\}}, U_{\{M\} \setminus \{x\}} | D_{\{x\}}, U_{\{x\}}) = P(D_{\{M\}}, U_{\{M\}}) / P(D_{\{x\}}, U_{\{x\}}),$$

where  $P(D_{\{M\}}, U_{\{M\}})$  and  $P(D_{\{x\}}, U_{\{x\}})$  are specified in formula (1), treating  $\{M\}$  and  $\{x\}$  as blocks, respectively. We note, however, that the marker set  $\{x\}$  is unknown a priori, and needs to be inferred jointly with other parameters in our Bayesian model.

### 3. Joint inference of diplotype blocks and disease association.

**3.1. The joint model.** To further incorporate epistatic interactions in formula (2), and to identify which SNPs are associated with the disease (response), we partition all SNPs (not blocks) into three groups as in BEAM [Zhang and Liu (2007)]. We introduce a latent  $L$ -dimensional indicator variable ( $I$ ) to represent the group memberships of the  $L$  markers. For each marker  $i$ ,  $I_i = 0, 1, 2$  denotes three possible group memberships. SNPs belonging to group-2 are assumed to be jointly



associated with the disease, that is, epistasis, which are modeled by two joint multinomial distributions on the diplotypes over all group-2 SNPs—one for cases and one for controls. SNPs belonging to group-1 are assumed to be marginally associated with the disease if they belong to different blocks, and are modeled by mutually independent multinomial distributions conditional on the case–control status and the block structure. If multiple group-1 SNPs fall into one block, we model their diplotypes jointly, that is, group-1 SNPs within blocks become dependent of each other. If there are both group-1 and group-2 SNPs within one block, we model the diplotypes of group-1 SNPs within the block conditional on the diplotypes of the group-2 SNPs within the block. SNPs belonging to group-0 are the remaining SNPs unrelated to the disease status. We again model the distribution of group-0 SNPs within a block by multinomial distributions, with common parameters for cases and controls. We further assume conditional independence of group-0 SNPs between blocks, conditional on the group-1 and group-2 SNPs. More precisely, within each block, we let  $\{x_2\}$  denote the set of group-2 SNPs, let  $\{x\}$  denote the union of the group-1 and group-2 SNPs, and let  $\{M\}$  denote all SNPs. We revise formula (2) to take the form of a conditional probability function:

$$(3) \quad \begin{aligned} &P(D_{\{M\} \setminus \{x_2\}}, U_{\{M\} \setminus \{x_2\}} | D_{\{x_2\}}, U_{\{x_2\}}) \\ &= \frac{P(D_{\{x\}})P(U_{\{x\}})P(D_{\{M\} \setminus \{x\}}, U_{\{M\} \setminus \{x\}} | D_{\{x\}}, U_{\{x\}})}{P(D_{\{x_2\}})P(U_{\{x_2\}})}. \end{aligned}$$

Thus, group-0 and group-1 SNPs are no longer mutually independent as in BEAM1, but are related to each other via the block structure. With epistasis considered, the mutual independence between blocks in model (2) becomes conditional independence given group-2 SNPs. For notational simplicity, we omit variable  $I$  in (3), but both  $\{x\}$  and  $\{x_2\}$  are determined by  $I$ .

Given a particular block partition  $B$  and SNP group memberships  $I$ , we express the joint probability function of the entire case–control data as

$$(4) \quad \begin{aligned} P(D, U | B, I) &= P(D_2 | B, I) P(U_2 | B, I) \\ &\times \prod_{\{M\}=[a,b] \in B} P(D_{\{M\} \setminus \{x_2\}}, U_{\{M\} \setminus \{x_2\}} | D_{\{x_2\}}, U_{\{x_2\}}, B, I), \end{aligned}$$

where  $D_2$  and  $U_2$  denote the case and control genotypes of group-2 SNPs, respectively, and the product term is defined in formula (3).

**3.2. Choice of prior distributions.** We set the prior distribution of the block variable  $B$  as the product of independent Bernoulli probabilities  $P(B) = p^{|B|}(1-p)^{L-|B|}$ , where  $|B|$  denotes the sum of indicators in  $B$ . According to the block distributions estimated in European and Asian populations by Gabriel et al. (2002), we assume that there are 50,000 blocks in the human genome a priori, and thus we set  $p = \min(0.5, 50,000R/(3 \times 10^9L))$ . Here,  $R$  denotes the length of the region



spanned by the  $L$  SNPs, and  $3 \times 10^9$  is the length of the human genome. A smaller value of  $p$  will help the method identify larger blocks, and a larger  $p$  will tend to identify smaller blocks. As the sample size (number of individuals) increases, however, the impact of the prior choices diminishes quickly. To avoid overfitting the blocks, we further impose a restriction that the maximum number of observed distinct diplotypes in a block must be smaller than  $(N_d + N_u)/10$ .

We set the prior distribution of the SNP membership variable  $I$  as a product of independent multinomial distributions,  $P(I) = \prod_{i=0}^2 p_i^{|I_j=I|}$ , where  $\{p_0, p_1, p_2\}$  denote the prior probability of each SNP belonging to group 0, 1, and 2, respectively. By default, we set  $p_1 = p_2 = \min(0.1, 5/L)$ , and  $p_0 = 1 - p_1 - p_2$ . That is, we assume there are 10 SNPs associated with the disease a priori, where 5 are marginally associated with the disease, and 5 are associated through epistasis. Our choice of the prior reflects that there are just a few SNPs truly associated with the disease in a GWA study (where many other significant SNPs are due to LD effects). Increasing this prior (and also increasing the significance level) in the BEAM2 program may help identify additional SNPs of moderate to low effects. To avoid overfitting in interaction mapping, we further set an upper bound to the order of interactions by  $\ln_3((N_d + N_u)/10)$ . For example, when the sample size is 1,000, our method can detect up to 4-way interactions. Overall, changing the values of  $p_1, p_2$  may affect the posterior distribution of SNPs in groups 1 and 2, but the effects will diminish as the sample size increases.

Finally, the joint model of the observed genotype data in cases ( $D$ ) and controls ( $U$ ), the block variable ( $B$ ), and the SNP membership ( $I$ ), is written as

$$(5) \quad P(D, U, B, I) = P(D, U|B, I)P(B)P(I),$$

where the conditional distribution of  $(D, U)$  given  $(B, I)$  is specified in formula (4).

**3.3. MCMC updates.** The parameters of interest in our model are the block partition  $B$  and SNP membership  $I$ . We develop Metropolis–Hastings (MH) algorithms [Liu (2001)] to update  $B$ , and, simultaneously, we develop a mix of a Gibbs sampler and Metropolis–Hastings algorithm to update  $I$ . The posterior distribution of  $(B, I)$  is then output for further analysis.

To explore all possible block partitions, we propose the following MH-moves: given a current block configuration  $B$ , we randomly select a block and

- (1) divide the block into two new blocks at a random position;
- (2) merge two adjacent blocks into one block; and
- (3) randomly shift a block boundary to either left or right by  $k$  SNPs, where the shifting amount is constrained by other block boundaries.

The proposed move produces a new block partition  $B'$ , and the move is accepted with probability  $r = \min\{1, \frac{P(D, U, I, B')q(B \rightarrow B')}{P(D, U, I, B)q(B' \rightarrow B)}\}$ , where  $q(B \rightarrow B')$  denotes the

probability of updating from  $B$  to  $B'$ , and  $P(D, U, B, I)$  is calculated from the full model (5). In our implementation, we chose the three types of MH moves with probabilities 0.1, 0.1, and 0.8, respectively, and we require a block to contain at least one SNP.

To update the SNP membership variable  $I$ , we updated the membership  $I_i$  of SNP  $i$  by calculating the posterior distribution of  $I_i = 0, 1, 2$  given all other model parameters and the data. We also propose a MH-move to switch the group memberships of two SNPs and accept the move based on MH-ratios. Per MCMC iteration, we first run the Gibbs sampler to update the memberships of all SNPs once, and then we run the MH-sampler to switch each SNP in group-1 and group-2 once with SNPs in other groups.

#### 4. Follow-up tests and generalization of the method.

4.1. *A test of significance based on the Bayes factor.* Although inference can be directly made from the posterior probabilities output by BEAM2, the users may want to further evaluate the statistical significance of the results in a frequentist way. In BEAM [Zhang and Liu (2007)], we developed a novel Bayes factor, called B-stat, to evaluate whether a SNP or a set of SNPs are significantly associated with the disease, where the SNP set is selected by BEAM2 in our case.

For a set of  $M$  SNPs to be tested, the null hypothesis is that all  $M$  SNPs are not associated with the disease. Here,  $M = 1, 2, 3, \dots$  represents single-SNP, 2-way, and 3-way interactions, etc. B-stat for the set of  $M$  SNPs is defined as

$$(6) \quad B_M = \ln \frac{P_A(D_M, U_M)}{P_0(D_M, U_M)} = \ln \frac{P(D_M)[P(U_M) + \prod_{j \in M} P(U_j)]}{P(D_M, U_M) + \prod_{j \in M} P(D_j, U_j)}.$$

Here,  $P_0(D_M, U_M)$  denotes the null genotype distribution (i.e., no disease association) at the  $M$  SNPs in cases and controls, and  $P_A(D_M, U_M)$  denotes the alternative genotype distribution (disease association). Under the null model, we assume that the genotypes in both cases and controls follow the same distribution, whereas under the alternative model, they follow different distributions. We choose both  $P_0(D_M, U_M)$  and  $P_A(D_M, U_M)$  as an equal mixture of two distributions: one that assumes independence among the  $M$  SNPs in controls (and also in cases under the null model), which yields the product terms in formula (6), and the other that assumes a saturated joint distribution of all the  $M$  SNPs. Note that the form of each term in formula (6) is defined in formula (1).

An interesting feature of B-stat is that it uses a mixture model to accommodate the possibility that the  $M$  SNPs may or may not be in linkage equilibrium (independence). As a result, using B-stat will be more powerful than using a standard likelihood ratio test or a chi-square test of associations when the  $M$  SNPs under testing are in LD in controls.

We have previously shown that, under the null hypothesis of no disease association, B-stat follows asymptotically a shifted chi-square distribution with  $3^k - 1$

degrees of freedom [Zhang and Liu (2007)]. The shifting parameter can be computed explicitly, which is determined by the sample size ( $N_d, N_u$ ), the interaction size  $M$ , and the Dirichlet hyper-parameter  $\{\alpha_h\}$ . Briefly speaking, the shifting parameter is proportional to  $-(3^M - 1) \ln(N_d N_u / (N_d + N_u))$ , and, thus, the larger the number of individuals collected, or the more SNPs involved in an interaction, the smaller the shifting parameter will be. In addition, if large hyper-parameters  $\{\alpha_h\}$  for the diplotype frequency parameters are used, the shifting parameter will be large too. Note that we want the B-stat to be small (e.g.,  $<0$ ) when the  $M$  SNPs are not associated with the disease, and, hence, the users should use small values for  $\{\alpha_h\}$ , such as the default values we used in our model.

**4.2. Generalization to classification problems with discrete covariates.** Let  $\mathbf{Y}$  be the  $n \times 1$  binary response vector, and let  $\mathbf{X} = (\mathbf{X}_1, \dots, \mathbf{X}_p)$  be the  $n \times p$  covariates matrix, with each covariate  $\mathbf{X}_j$  taking on  $k_j$  discrete (ordinal or categorical) values. The standard case-control genetic study setting can be viewed as using response variables  $\mathbf{Y}$  (i.e., case-control status) to fish out relevant predictors  $\mathbf{X}_j$  (i.e., SNPs). The epistasis mapping methods BEAM attempt to find those  $\mathbf{X}$ 's that interactively affect  $\mathbf{Y}$ . Both BEAM and the block-based method BEAM2 can be easily extended to infer a classification model.

The idea of both BEAM [Zhang and Liu (2007)] and BEAM2 is to partition the  $p$  covariates in  $\mathbf{X}$  into three nonoverlapping groups, such that one group contains covariates unrelated with  $\mathbf{Y}$ , and the other groups contain covariates either independently or jointly related with  $\mathbf{Y}$ . The partition of the covariates is an unobserved latent structure. Given a particular group partition of the covariates  $I$ , we can compute  $P(\mathbf{X}|\mathbf{Y}, I)$  as in Zhang and Liu (2007), which is analogous to that in a naïve Bayes model. BEAM2 further segments the covariates  $\mathbf{X}$  into “blocks” of highly correlated variables, and treats blocks as mutually independent. This is achieved by introducing a block indicator variable  $B$ , which is updated iteratively together with the variable selection indicator  $I$ .

To predict the classification of a new observation with covariates  $\mathbf{x}_{new}$  based on the training data  $(\mathbf{X}, \mathbf{Y})$ , we compute  $P(\mathbf{X}, \mathbf{x}_{new}|\mathbf{Y}, \mathbf{y}_{new} = 1)$  and  $P(\mathbf{X}, \mathbf{x}_{new}|\mathbf{Y}, \mathbf{y}_{new} = 0)$ , respectively, using the BEAM (or BEAM2) algorithm, and obtain the odds ratio

$$P(\mathbf{x}_{new}|\mathbf{X}, \mathbf{Y}, \mathbf{y}_{new} = 1) / P(\mathbf{x}_{new}|\mathbf{X}, \mathbf{Y}, \mathbf{y}_{new} = 0).$$

The prior  $P(\mathbf{y}_{new} = 1)$  can be estimated from the prior knowledge of class distribution, such as the prevalence of a particular disease in the population, which then leads to the posterior predictive probability for  $\mathbf{y}_{new} = 1$ . A computationally more attractive way to do the computation is to output the latent variable partition and block partition structures ( $I$  and  $B$  in our case) from their joint posterior distribution inferred by the MCMC procedure of BEAM, and then average the conditional odds over all the sampled  $I$  and  $B$ :

$$P(\mathbf{x}_{new}|\mathbf{X}, \mathbf{Y}, I, B, \mathbf{y}_{new} = 1) / P(\mathbf{x}_{new}|\mathbf{X}, \mathbf{Y}, I, B, \mathbf{y}_{new} = 0).$$

We tested this latter approach in a preliminary study and found the results quite satisfactory.

The effect of BEAM2 is somewhat analogous to that of *elastic net* [Zou and Hastie (2005)] and *group lasso* [Yuan and Lin (2006)]. All methods attempt to address the phenomena that groups of covariates tend to demonstrate associations with the response together, and within groups the covariates are highly correlated. Different from *elastic net* and *group lasso*, BEAM2 infers the covariate groups and also the informative covariates within groups jointly in a coherent probability framework. As a consequence, BEAM2 allows sparse variable selection at both the group level and the individual variable level within groups, whereas *elastic net* and *group lasso* do sparse selection only at the group level. In a recent technical report, Friedman, Hastie and Tibshirani (2010) attempt to achieve a similar sparse selection effect as BEAM2 (sparse selection at both group and individual levels) by introducing an additional penalty term.

Other important distinctions between BEAM2 (or BEAM) and those lasso-based methods are the following: (a) the use of the naïve Bayes framework to model  $\mathbf{X}$  given  $Y$  to greatly alleviate the overfitting problem; (b) the ability to incorporate interaction terms without incurring a huge computational burden (with MCMC iterations); and (c) the adoption of the Bayesian variable selection principle, which is equivalent to using a more desirable  $L_0$  penalty. The cost of these advantages is that both BEAM and BEAM2 have to compute via MCMC without a guarantee of always finding the optimal solution. Empirically, however, the computational speed of BEAM and BEAM2 is no worse than that required by lasso-type algorithms when the number of covariates is large.

## 5. Simulation studies and algorithm comparisons.

**5.1. Block partition of HLA data.** We first used the HLA region on human chromosome 6 to evaluate the block partitions inferred by our method. The HLA region is one of the few regions in the human genome in which recombination hotspots have been experimentally verified [Jeffreys, Ritchie and Neumann (2000); Jeffreys, Kauppi and Neumann (2001)]. We downloaded the genotype data of 50 unrelated UK Caucasian semen donors from Jeffreys AJ's website. The data covers a 216 kb region with 296 genotyped biallelic markers spanning from the upstream of gene *HLA-DNA* to gene *TAP2* in the MHC Class II region. It is known that this region contains several prominent recombination hotspots [Jeffreys, Ritchie and Neumann (2000); Jeffreys, Kauppi and Neumann (2001)]. We therefore examined the relationship between the experimentally verified recombination hotspots and the SNP-block boundaries inferred by BEAM2. We used both haplotypes and genotypes to evaluate our method, where the HLA haplotypes were first inferred by CHB [Zhang, Niu and Liu (2006)] from the genotype data. We also simulated 1,000 individuals from the inferred HLA haplotypes using HAPGEN [Marchini et al. (2007)] to evaluate the performance of BEAM2 with a larger sample

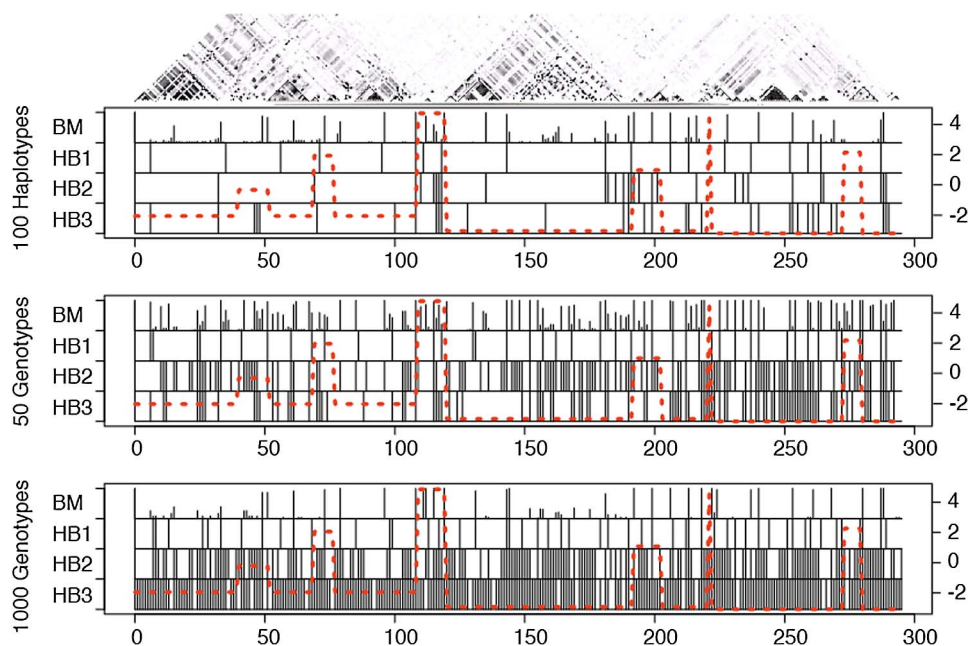


FIG. 1. Block partition results on the HLA data. The top panel shows the pairwise SNP LD calculated from 100 HLA haplotypes by Haploview. The second, third, and forth panels show the block partition results by BEAM2 (BM) and HapBlock (HB1, HB2, HB3) on three HLA data sets, respectively. HB1 uses rule 1: common haplotypes; HB2 uses rule 2: pairwise  $D'$ ; and HB3 uses rule 3: four-gamete test. Experimentally verified recombination rates are shown as dashed red lines (the unit cM/Mb in the natural logarithm scale is shown on the right).

size. As a comparison, we applied HapBlock [Zhang et al. (2002b)], a dynamic programming-based algorithm of block partitioning, to the same sets of data.

With 100 haplotypes inferred from the 50 individuals, BEAM2 produced accurate block partitions that correspond well to the visual blocks displayed by Haploview [Barrett et al. (2005)]. The block boundaries also coincide with the known recombination hotspots within the HLA region (Figure 1). It is further observed that, for the haplotype data, the blocks inferred by BEAM2 are very similar to those obtained by HapBlock. Unlike our model-based method, HapBlock requires the user to specify ad hoc block partition rules, which can result in undesirable partitions. We used three different rules to define blocks: (1) common haplotypes, defined as a haplotype  $>5\%$  in the sample, cover 80% of samples in a block; (2) at least 80% SNP pairs with  $D' > 0.5$  in a block; or (3) four-gamete test on common haplotypes ( $>5\%$ ) in a block. The blocks partitioned by HapBlock using each rule are also shown in Figure 1.

Using the genotype data of the 50 individuals, we obtained very different results between BEAM2 and HapBlock, and between the three different rules of block partitions. Except for the first rule of HapBlock, all other methods produced

TABLE 1  
*Number of blocks inferred in HLA data sets*

Data	BEAM2	BEAM2-10p	HapBlock-1	HapBlock-2	HapBlock-3
100 haplotypes	32	39	20	30	34
50 genotypes	81	87	46	147	106
1,000 genotypes	35	36	56	184	268

“BEAM2-10p” denotes BEAM2 applying a 10 times larger prior probability than the default prior on the block boundary variable. HapBlock-1, 2, 3 denotes HapBlock applying three different block partition rules.

a large number of small blocks. Small blocks generated by BEAM2 are due to the small sample size of 50 individuals, based on which the correlation between SNPs is hard to detect using our likelihood model. The posterior probabilities of block boundaries output by BEAM2, however, can be used as a measure of uncertainty in block partitions. In comparison, HapBlock and many other block partitioning methods only provide a single partition solution without measuring block uncertainty.

Using the simulated genotypes of 1,000 individuals, we again observed very different results shown in Figure 1. BEAM2 produced the cleanest block partitions that corresponded well to the visual block boundaries and to the known recombination hotspots. The  $D'$  rule and the four-gamete test rule via HapBlock again failed to produce reasonable partitions, of which most blocks were singletons.

We further show in Table 1 the number of blocks inferred by each method in the three data sets. We observed that BEAM2 performed well (by which we roughly mean that the number of estimated blocks is small, as is true in the HLA region) in both the haplotype data and the 1,000 individuals’ genotype data. Because modeling genotypes (diplotypes) requires a much larger set of parameters than modeling haplotypes, BEAM2 is expected to perform worse in the 50 individuals’ genotype data. As the number of individuals increased to 1,000, however, our model-based approach produced very similar partitions as that obtained in the haplotype data. In comparison, HapBlock only preformed reasonably well in the haplotype data, but produced many small blocks and singletons in the other two data sets for all three block partition rules applied. HapBlock performed the worst in the 1,000 individuals’ genotype data, indicating that the ad hoc rules applied by HapBlock do not produce consistent block partitions as sample size increases. We further show in Table 1 additional results by BEAM2 using a 10 times larger prior on the block boundary variable, that is, we expect 10 times more blocks a priori. We observed that the estimated posterior number of blocks did not increase much, particularly in the 1,000 individuals’ genotype data, indicating that BEAM2 is insensitive to the prior choice of block boundary variables. We also ran multiple MCMC chains to ensure proper convergence.



*5.2. Simulation study using HapMap data.* To mimic real genetic data observed in human populations, we first randomly select a region in the human genome that contains 1,000 Illumina HapMap 300k tagSNPs. The region also contains about the same number of additional SNPs from HapMap PhaseII tagged by these tagSNPs, which we refer to as nontagging SNPs. Two nontagging SNPs in the region are randomly selected as disease SNPs. Given a disease model, we set the marginal effect size (log odds ratio minus 1) per disease SNP at 0.5 and choose a disease minor allele frequency (MAF) per locus from (0.05, 0.10, 0.20, 0.50). Given a marginal effect size and a choice of MAF, we then calculate the diplotype frequencies over the disease SNPs in cases and controls, respectively [this is similarly done as presented in Zhang and Liu (2007)]. According to the case-control diplotype frequencies over the disease SNPs, we randomly sample 1,000 cases and 1,000 controls from a pool of individuals without replacement. The pool consists of 10,000 control individuals generated by HAPGEN [Marchini et al. (2007)] using HapMap European sample (parents only) at odds ratio = 1, that is, no disease association. Our simulation procedure is more economical than a direct approach that generates one individual at a time and determines its disease status conditional on the disease genotypes and penetrance, because the direct approach may generate many more controls before obtaining enough cases. Finally, we remove all nontagging SNPs from the data including the two disease SNPs (which are typically unobserved in a GWA study), and obtain a case-control data set containing 1,000 Illumina HapMap tagSNPs.

To evaluate the association mapping performance of our method, we simulated case-control data sets based on the HapMap sample under three disease models shown in Table 2. Each disease model assumes 2 loci in the genome contributing to the disease risk. While the first model assumes no interactions, the other two models assume different types of interactions. Using the simulated data sets, we compared the performance of BEAM2 to BEAM1. We also implemented a third method that maps associations and interactions based on predetermined block structures. This third method serves as an intermediate method between BEAM1, which is not block-based, and BEAM2, which infers block structures and maps associations simultaneously. We used three different levels of parameters (from stringent to liberal) to define blocks using existing software, and we treated the diplotypes within each inferred block as genetic alleles. The third method using three different block definitions are hereafter referred to as Block1, Block2, and Block3, respectively [more details of the third method can be found in the Supplementary Material, Zhang, Zhang and Liu (2011)]. To compare the performance of all methods, we ranked SNPs according to the association posterior probabilities output by each method estimated for each data set. We then calculated how often a method ranked the disease related SNPs among the top SNPs. A SNP is regarded as being correctly identified as disease related if it is within 5 SNPs on either side of a true disease locus.



TABLE 2  
*Disease models used in simulation study*

Risk	<i>A/A</i>	<i>A/a</i>	<i>a/a</i>
Model 1			
<i>B/B</i>	1	$1 + \theta$	$(1 + \theta)^2$
<i>B/b</i>	$1 + \theta$	$(1 + \theta)^2$	$(1 + \theta)^3$
<i>b/b</i>	$(1 + \theta)^2$	$(1 + \theta)^3$	$(1 + \theta)^4$
Model 2			
<i>B/B</i>	1	1	1
<i>B/b</i>	1	$(1 + \theta)^2$	$(1 + \theta)^3$
<i>b/b</i>	1	$(1 + \theta)^3$	$(1 + \theta)^4$
Model 3			
<i>B/B</i>	1	1	1
<i>B/b</i>	1	$1 + \theta$	$1 + \theta$
<i>b/b</i>	1	$1 + \theta$	$1 + \theta$

Each table cell lists the relative risk of the corresponding genotype combination. Genotypes with risks equal to 1 have no effects to the disease. The parameter  $\theta$  is computed according to the specified marginal effects (0.5 in our simulation) and disease MAFs (0.05, 0.1, 0.2, 0.5).

As shown in Figure 2, under all parameter settings, BEAM2 performed the best among all tested methods, where Block1, Block2, Block3, and BEAM1 all performed similarly. When disease allele frequency was low ( $f = 0.05$ ), the power curves of all methods looked similar, but a closer inspection of the top 5 ranked SNPs showed that BEAM1 only had  $\sim 50\%$  chance to capture disease related SNPs relative to BEAM2. When disease alleles were common in the population ( $f = 0.10, 0.20, 0.50$ ), the advantage of the BEAM2 model becomes obvious. Comparing the power curves for Model 2 and Model 3, we observed that the power of BEAM2 increased much faster than that of BEAM1 among the top 2 or 3 SNPs. We did not observe this behavior in Model 1, which has no interactions. It thus indicates that using SNP-blocks can increase the power of mapping both single SNP and multi-SNP interaction associations. All methods compared here are Bayesian methods that output posterior probabilities of disease associations. It therefore indicates that our treatment of LD in BEAM2 is more appropriate than using either predetermined blocks or a Markov chain model (as in BEAM1).

To further declare statistical significance, existing significance estimation methods adjusting for multiple comparisons should be used, such as the Bonferroni correction applied to B-stat introduced in BEAM1 [Zhang and Liu (2007)]. We compared BEAM2 with single SNP chi-square tests using the above disease models [Supplementary Table S1, Zhang, Zhang and Liu (2011)], and observed that BEAM2 performed better than the chi-square test for interaction Model 2 and Model 3, but performed the same for the noninteractive Model 1.

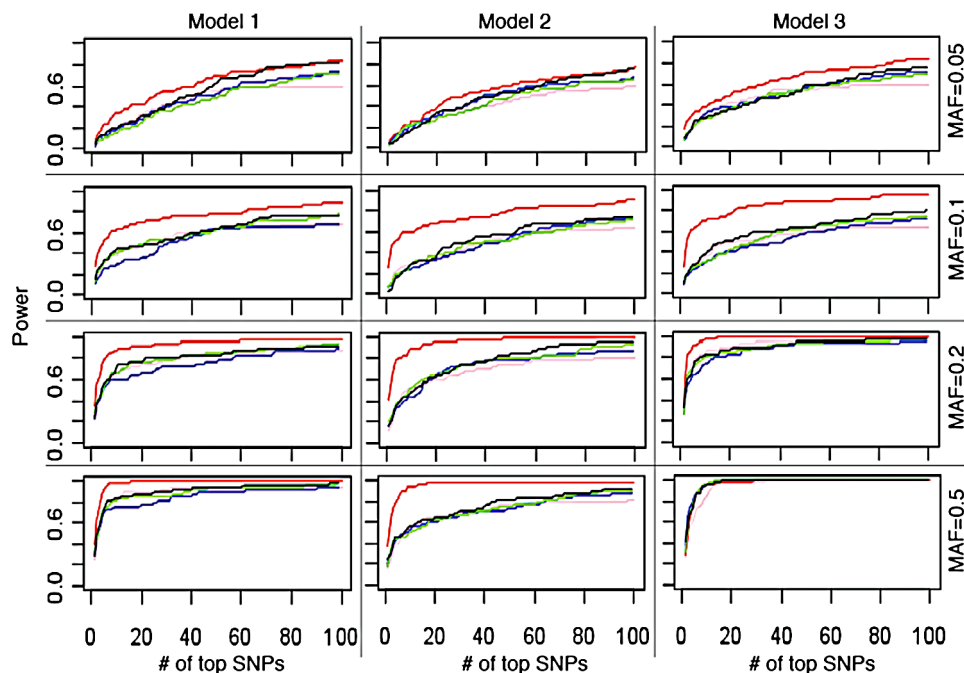


FIG. 2. Power comparison for BEAM1, BEAM2, Block1, Block2, and Block3 using simulated data from the HapMap data. Under each simulation setting and from 50 data sets, power (y-axis) is calculated as the proportion of disease-associated SNPs (within 5 SNPs of true disease loci) among top  $m$  SNPs (x-axis), ranked by the posterior probability of association. Each data set contains 1,000 candidate SNPs in 1,000 cases and 1,000 controls. The disease allele frequency is 0.05, 0.10, 0.20, and 0.50, respectively. The marginal effect size of each disease SNP (unobserved) is 0.5. Pink: BEAM1; Red: BEAM2; Blue: Block1; Green: Block2; Black: Block3.

We also checked the performance of our MCMC sampling algorithm. As shown in Figure 3, using a simulated data set from disease Model 2, the lag of autocorrelation of our Markov chain is short, indicating fast convergence of the Markov chain. We further compared the posterior distribution of SNP associations from 4 independent runs of BEAM2, and we observed close agreement between runs. In practice, the Markov chain could converge to local modes, particularly if the data contain many SNPs with complicated block structures. If block structures are of primary interest, we suggest running BEAM2 in several runs to check if the block partition results obtained in different runs are consistent. More advanced MCMC algorithms, such as parallel tempering [Liu (2001)], could further alleviate the local mode problems in MCMC sampling.

Using BEAM2, we can estimate the number of disease associated SNPs around a disease locus by the sum of posterior probabilities of associations over all SNPs within a neighborhood of a candidate locus. Given our block-based association model, the number of associated SNPs does not include SNPs whose disease as-

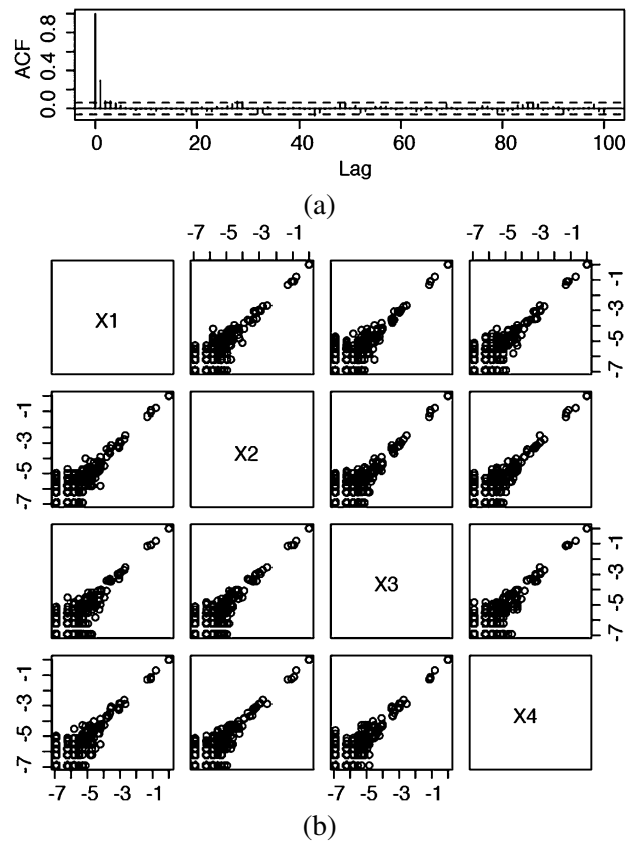


FIG. 3. Performance of MCMC. (a) Autocorrelation plot obtained by running BEAM2 on a simulated data set of 1,000 cases, 1,000 controls, and 1,000 SNPs under disease Model 2. A total of 1,000 iterations after burn-in are used to calculate the plot. (b) Posterior probabilities (in logarithm scale) of disease association (marginal and epistatic) per SNP compared across 4 independent runs of BEAM2.

sociation is merely created by LD, and, hence, our estimates are more appropriate than a naïve count of significant SNPs within the neighborhood. As shown in Figure 4, around a 100-kb neighborhood of every simulated disease locus in disease Model 1, the estimated number of disease associated SNPs by BEAM2 is around 1 when the association signal is sufficiently strong, even if there are many significant SNPs in the neighborhood. The extra significant SNPs created by LD make the localization of disease locus difficult. This result highlights the importance of BEAM2 that performs automatic variable selection within blocks. Rather than reporting diluted small posterior probability of association over many neighboring SNPs in LD, BEAM2 was able to select the strongest contributing SNP within blocks (with large posterior probabilities of association) in our simulation study.

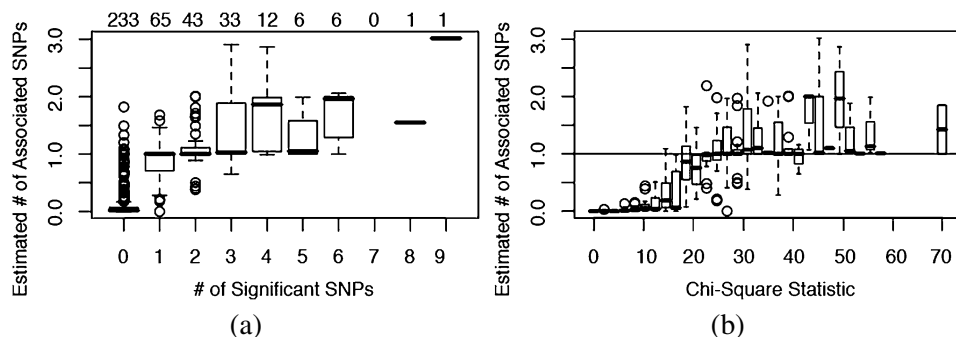


FIG. 4. Estimated number of disease associated SNPs (y-axis) plotted against (a) the number of significant SNPs ( $>0.05$  family-wise) and (b) the maximum single SNP chi-square statistics (x-axis) within a 100 kb neighborhood per disease locus. In (a), the number of loci in each box plot is further shown on the top. The plots are computed from 200 simulated data sets of disease Model 1.

**6. Application to WTCCC type 1 diabetes data.** We applied BEAM2 to analyze the T1D data set from the WTCCC project [WTCCC (2007)]. The data set contains 2,000 T1D patients, 1,504 controls from 1958 Birth Cohort (58C), and 1,500 additional controls from the National Blood Service (NBS). Given our limited computation resources (computation time, which would require several days to analyze half a million SNPs in this data set; and memory usage, which would require  $>4$  Gb for half million SNPs), we applied BEAM2 to the top 10% SNPs ranked by marginal associations with T1D on all autosomes. We further filtered out SNPs with bad clustering, SNPs violating Hardy–Weinberg equilibrium in controls at  $10^{-5}$  significance, and “almost nonpolymorphic” SNPs of which  $>95\%$  samples have the same genotypes. The final data set contained 42,470 SNPs in 5,004 individuals. We ran BEAM2 with 5 independent MCMC chains. Figure 5 shows the averaged posterior probabilities of T1D associations. Note that selecting top 10% SNPs does not imply sparse and less correlated SNPs, because SNPs in LD tend to be in or out of the top 10% list together. Two alternative ways to reduce the number of SNPs are to run BEAM2 on each individual chromosome for chromosome-wise epistasis, or on previously detected disease associated regions.

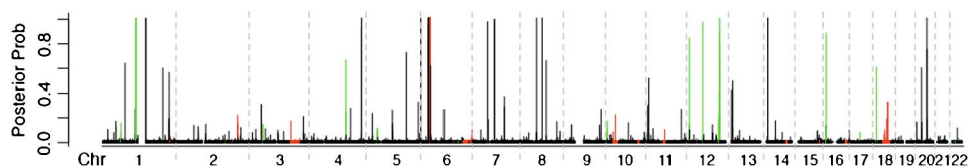


FIG. 5. SNP-wise posterior probabilities (y-axis) of T1D associations in 22 autosomal chromosomes (x-axis, chromosomes are separated by grey dashed lines). Previously reported T1D associated genes are highlighted in green, and candidate T1D associated genes, as defined at T1Dbase, are highlighted in red.

6.1. *Result summary and highlights.* We first compared our results with the SNPs reported in the original WTCCC paper (2007). As expected, we found that the SNPs reported by BEAM2 and by the original WTCCC analysis are highly consistent. All strongly associated SNPs reported in WTCCC are significant in our analysis, and all strongly and moderately associated SNPs reported in WTCCC have posterior probability  $> 0.1$  by BEAM2 [Supplementary Table S2, Zhang, Zhang and Liu (2011)]. We further compared our results with known T1D associations obtained from T1Dbase ([www.t1dbase.org](http://www.t1dbase.org)). Among the 55 SNPs (or cluster of SNPs) output by BEAM2 with posterior probabilities greater than 0.1, 17 (31%) overlapped with known T1D associated regions, including some well-known genes such as *PTPN22* (1p13), *CTLA4* (2q33), MHC (6p21), and *IL2RA* (10p15).

In addition to the previously reported T1D genes, BEAM2 reported some novel T1D associated loci. A list of likely T1D associations detected by BEAM2, for both single SNPs (if  $p$ -value  $< 5e-7$ ) and two-way joint associations (if  $p$ -value  $< 5e-10$ ), is shown in Tables 3 and 4, respectively. For example, we detected 7 loci, among which two SNPs in short distance form strong joint associations with T1D ( $p$ -value  $< 5e-10$ ). These loci are not identifiable using single SNP tests, but captured by BEAM2 as multi-SNP associations. We also found some likely long-distance and cross-chromosomal interaction associations with T1D ( $p$ -value  $< 5e-11$ ). One example is the joint association between SNP rs3132676 in the classic MHC region on chromosome 6 and SNP rs9376523, which is 111 Mb away on the same chromosome. This SNP pair is likely interacting because their genotypes are strongly correlated in cases (nominal  $p$ -value

TABLE 3  
*Strong single SNP associations with T1D*

SNP	Position	$p$ -value	T1Dbase	Gene
rs6679677*	chr1: 113.8–114.2 Mb	0	Yes	<i>PTPN22</i>
rs9405484	chr6: 1.4 Mb	1.33e–8	No	<i>FOXC1</i>
MHC*	chr6: 25–35 Mb	0	Yes	<i>MHC</i>
rs6592988	chr7: 52.1 Mb	2.87e–7	Yes	<i>COBL</i>
rs11984645	chr8: 55.2 Mb	7.06e–11	No	<i>MRPL15</i>
rs11782342	chr8: 73.9 Mb	5.70e–12	No	<i>KCNB2</i>
rs11052552	chr12: 9.7 Mb	2.61e–7	Yes	<i>CLEC2D</i>
rs11171739*	chr12: 54.8 Mb	2.35e–11	Yes	<i>ERBB3</i>
rs17696736*	chr12: 109.8–110.9 Mb	0	Yes	<i>CCDC63, NAP1</i>
rs12924729*	chr16: 11.1 Mb	1.01e–7	Yes	<i>CLEC16A</i>

SNPs showing strong associations ( $p$ -value  $< 5e-7$ ) with T1D by single SNP test.  $p$ -value: nominal  $p$ -value of associations. T1Dbase: whether the locus is documented in T1Dbase. Gene: nearest gene.

\*Additional SNPs in its neighborhood also show strong marginal associations.

TABLE 4  
Strong two-SNP associations with T1D

SNP1	Pos1	Pval1	Gene	SNP2	Pos2	Pval2	Gene	JointP
rs7525703	chr1: 143	2e-2	<i>PRKAB2</i>	rs2077749	chr1: 143	3e-6	<i>PRKAB2</i>	2e-12
rs6809441	chr3: 41	4e-2	<i>ULK4</i>	N/A	chr3: 41	7e-2	<i>ULK4</i>	5e-11
rs330483	chr4: 176	4e-6	<i>ADAM29</i>	rs330504	chr4: 176	1e-1	<i>ADAM29</i>	2e-11
rs6906469	chr6: 10	1e-3	<i>DFCCI</i>	rs659964*	chr12: 111	2e-6	<i>ACAD10</i>	4e-11
rs3132676*	chr6: 30	1e-5	<i>TRIM40</i>	rs9376523	chr6: 141	6e-4	<i>TXLNB</i>	3e-11
rs9296661	chr6: 52	4e-4	<i>PKHDI</i>	rs1265566*	chr12: 110	1e-6	<i>CUTL2</i>	5e-11
rs13340508	chr7: 75	9e-3	<i>CCL24</i>	rs17361077	chr7: 75	4e-5	<i>CCL24</i>	6e-12
rs4838140	chr9: 124	2e-5	<i>NEK6</i>	rs7860360	chr9: 125	6e-4	<i>SCAI</i>	4e-10
rs11104868	chr12: 87	1e-4	<i>KITLG</i>	rs7961663*	chr12: 110	3e-6	<i>CUTL2</i>	1e-11
rs1958305	chr14: 23	5e-5	<i>DHRS2</i>	rs12100601	chr14: 23	2e-2	<i>DHRS2</i>	0
rs7262414	chr20: 40	2e-5	<i>PTPRT</i>	rs2867064	chr20: 40	7e-2	<i>PTPRT</i>	0

Pairs of SNPs showing strong joint T1D associations ( $p$ -value  $< 5e-10$ ) by B-stat. If multiple SNP pairs are located around the same loci, only one pair is shown. Pval1, Pval2, JointP represent nominal  $p$ -values of SNP1, SNP2, and their joint associations, respectively.

\*SNP lies in known regions in T1Dbase.

8e-6 by test of independence), but not in controls (nominal  $p$ -value 0.91). Although most two-way associations did not pass the Bonferroni adjusted significance level in the genome scale, the short-distance two-way associations are significant if only considering local joint associations.

We further examined possible confounding effects of population structures in the T1D data using a logistic regression model. The regional information of WTCCC individuals is included as dummy covariates. We observed that the test statistics of the detected SNP associations remained almost unchanged before and after the adjustment of population origins. We further randomly selected 10,000 SNPs genome-wide and compared the distribution of their association statistics with a chi-square distribution. The two distributions agreed well [see Supplementary Figure S1, Zhang, Zhang and Liu (2011)]. We therefore believe that population structure does not incur false positive associations in the WTCCC T1D data.

We finally checked the block partition results of BEAM2 on the T1D data. Given the large number of SNPs, we cannot visually inspect the blocks as we did for the HLA data. Alternatively, we computed the genetic distance between adjacent SNP pairs in the T1D data, using a genetic map constructed from the HapMap CEU sample. Then, we checked how frequent a block boundary is inferred between SNP pairs in certain genetic distance. Intuitively, the more distant two SNPs are located, the more likely a block boundary occurs. As shown in Figure 6, our method inferred almost 100% block boundaries between SNP pairs with genetic distance  $> 1$  cM, and the frequency of block boundary decreases as the SNP pairs get closer. The block partitions between the 5 runs of BEAM2 are consistent, with an average correlation coefficient of 0.96.

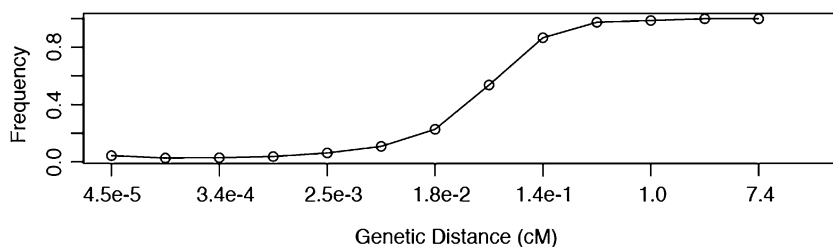


FIG. 6. Frequency (y-axis) of block boundaries between adjacent SNP pairs of certain genetic distance (x-axis), calculated from the T1D data. The genetic distance is obtained from the HapMap CEU sample.

**6.2. Joint association patterns in the MHC region.** T1D is an autoimmune disease and genes in the MHC region play an important role in the immune system and autoimmunity [Nejentsev et al. (2007); Steenkiste et al. (2007)]. BEAM2 found a large number of SNPs within the MHC region showing extremely strong association signals with T1D. Several multi-locus joint associations within the MHC region are also detected. We therefore examined more closely a 10-Mb MHC region, including the extended MHC region (25–32 Mb) and the classic MHC region (32–35 Mb).

Within this 10 Mb region, we observed that the SNP pairs associated with T1D are more often strongly correlated in cases than in controls [see Supplementary Figure S2, Zhang, Zhang and Liu (2011)]. The joint associations spanned from the classic MHC region to the extended MHC region over a distance as long as 6.5 Mb. It has been previously reported that haplotype blocks containing the most susceptible alleles *HLA-DRB1*\*03 and *HLA-DRB1*\*04 within the *HLA-DR-DQ* region may extend as long as 2 Mb [Nejentsev et al. (2007)] into the extended MHC region. It is thus arguable that the joint associations observed between the MHC class II region and the extended MHC region are due to extensive LD. We used a more traditional approach, logistic regression, to test two-way interactions among SNPs within MHC conditioning on the *HLA-DR-DQ* haplotypes. We first used CHB [Zhang, Niu and Liu (2006)] to infer haplotypes over *HLA-DR-DQ* genes (32.6–32.8 Mb). After collapsing rare haplotypes with frequencies lower than 0.1% into one group and obtaining 91 groups of haplotypes, we regressed the T1D status on the 91 haplotype groups using 90 dummy variables, which resulted in 78 significant haplotype groups over the *HLA-DR-DQ* genes explaining most of the MHC class II associations with T1D (the 13 insignificant haplotype groups removed from the model only changed the model deviance by 9.7).

Conditioning on the 78 haplotype groups, we tested both main and interaction effects between pairs of SNPs, one in the nonclass II region (class I, class III, and the extended MHC region) and one in the class II region. In particular, we regressed the T1D status on every pair of SNPs and the 78 haplotype groups. The association statistic is the change of deviance before and after including the two SNPs in the



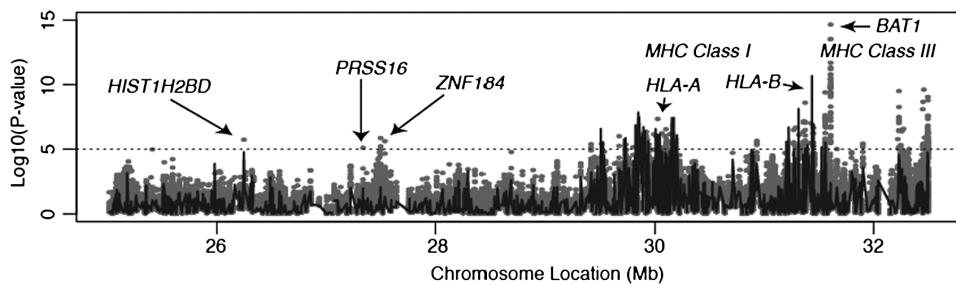


FIG. 7.  $\text{Log}_{10}$   $p$ -values (y-axis) of joint association test between SNPs in the nonclass II MHC region (x-axis) and SNPs in the MHC class II region (32.5–32.8 Mb), with main effects of the MHC class II SNPs subtracted and conditioned on 78 T1D associated haplotypes in HLA-DR-DQ. Each SNP has multiple  $p$ -values, shown as grey dots, corresponding to interactions with different MHC class II SNPs. The black line for the SNP indicates the  $-\log_{10}$   $p$ -value of the nonclass II SNP's main effect. A cutoff of  $10^{-5}$  for the  $p$ -values, shown as the dashed line, corresponds to roughly one false positive among all tests.

model. We further subtracted the main effect of the class II SNPs. As shown in Figure 7, we observed several peaks of  $p$ -values demonstrating significant main and interaction effects within the nonclass II MHC region. For example, we observed significant main effects of gene *HIST1H2BD* at 26.3 Mb (peak of black lines), interaction effects of gene *PRSS16* at 27.3 Mb and *ZNF184* at 27.5 Mb (peaks of grey dots), strong marginal effects of region 30–31.4 Mb including class I genes *HLA-A* and *HLA-B* (peaks of black lines), and strong interaction effects of gene *BAT1* at 31.6 Mb (peaks of grey dots). The associations of *PRSS16*, *ZNF184*, and *BAT1* have been previously reported [Nejentsev et al. (2007); Viken et al. (2009)], and our analysis further suggested that these genes are associated with T1D mainly through interactive effects with MHC class II genes, controlling the MHC class II haplotypes.

**7. Discussion.** In this article we proposed a model-based Bayesian method for simultaneous LD-block partitioning and multi-locus epistasis association mapping. Different from many block-based methods, we combined block partitioning and association mapping into a unified Bayesian model, where both block structures and SNP associations are iteratively learned through MCMC sampling. For block partitioning, our simulation study and real data analysis showed that BEAM2 produces accurate and consistent partitions of SNPs that compared favorably with other genetic knowledge than some existing methods. The posterior probabilities of block boundaries output by BEAM2 not only report the most likely block partitions but also measure the uncertainty of blocks. Compared to some existing methods using ad hoc partition rules, BEAM2 has two main advantages: (1) it provides soft partitions rather than hard partitions, that is, it provides a posterior distribution of block partitions given the data, rather than a single block partition result; soft partitions are necessary in regions with no dominant recombination hotspots

and block structures; and (2) it scales up well to large sample sizes and produces consistent results. In particular, we showed that our model-based method produces consistent block partitions as the number of individuals increases, whereas the other methods we tested produced drastically different results when more individuals from the same population are included. For association mapping, BEAM2 is more powerful than both the original BEAM algorithm, which accounts for LD using a Markov chain, and the methods using pre-estimated blocks. BEAM2 tests disease associations over uncertain block structures, where most SNPs around disease loci are filtered out, as their associations are created by LD with nearby disease SNPs.

The output of BEAM2 is a list of SNP-wise posterior probabilities of marginal and interaction associations. From a frequentist point of view, the identified SNPs can be further tested for genome-wide statistical significance. We previously introduced a Bayes factor-based statistics, B-stat [Zhang and Liu (2007)], for testing the significance of SNP associations. B-stat performs similarly to a 2-df chi-square test for single SNPs, but is more powerful for testing interactions of multiple SNPs. The same B-stat can be used to test the statistical significance of SNPs selected by the BEAM2 model, as we demonstrated in this paper.

When applied to the WTCCC T1D data, BEAM2 found all 5 statistically significant loci previously reported in Table 3 of WTCCC (2007), and captured 7 moderately (insignificant) associated loci listed in Table 4 of WTCCC (2007) with nontrivial posterior probabilities ( $>0.1$ ). BEAM2 further reported some novel two-way joint associations, including 7 SNP pairs ( $p$ -value  $< 5e-10$ ) in short-distance and 4 SNP pairs ( $p$ -value  $< 5e-11$ ) in long-distance. The local two-way associations indicate main effects of the related genes, which are, however, not detectable using single SNP tests alone. The long-distance two-way associations did not pass the genome-wide Bonferroni multiple testing control, but they may be justifiable as real interaction associations if a better multiple testing method is used and a replication study is performed. We further analyzed the well-known MHC region, and our analysis conditioning on the MHC class II haplotypes suggested the existence of interaction associations rather than MHC extended linkage alone. Given the complex nature and the important biological role of MHC in the human genome, our analysis is rather limited. Sophisticated analysis is in dire need to further reveal the genetic mechanisms underlying MHC and immune diseases.

The current BEAM model can be further improved in several ways. First, missing genotypes and unobserved SNPs in the case-control sample can be treated via imputation [Zhang (2011)]. Previous studies have shown that imputing untyped SNPs and missing genotypes from a reference panel can improve the power of disease association mapping [Marchini et al. (2007)]. Second, the current model only reports SNP-wise posterior probabilities of associations to the disease without providing a detailed analysis of association structures of the detected SNPs interactions. We have observed in the MHC region a bulk of SNPs exhibiting complex association structures with T1D. A post-analysis of the MHC region is thus

needed to delineate fine structures of the selected SNPs and interactions with respect to their disease effects and inter-relationships. Third, from a statistical point of view, it is critical to control the false-discovery rate. This is particularly important for multi-locus tests, which often involve a much larger number of simultaneous comparisons than single SNP tests. We are developing new statistical methods for evaluating genome-wide statistical significance of associations.

## SUPPLEMENTARY MATERIAL

**Additional Supporting Information and Results** (DOI: [10.1214/11-AOAS469SUPP](https://doi.org/10.1214/11-AOAS469SUPP); .doc). The file includes a naïve SNP-block model used in our comparison, verification of population structure in the sample, LD analysis of the MHC region, Chi-square results of our simulation study, and comparison of our results with previous results in the T1D WTCCC1 data.

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